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Assessing the effects of marine protected area (MPA) on a reef fish assemblage in a northwestern Mediterranean marine reserve: Identifying community-based indicators

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ABSTRACT

Marine protected areas (MPAs) are increasingly envisaged as a tool to manage coastal ecosystems and fisheries. Assessment of their performance with respect to management objectives is therefore important. A number of MPAs provided conservation benefits for fished species. Observed benefits do not apply to all species at all times, and responses to protection are also highly variable among fish taxa. Among the many empirical studies on marine reserves, only a few designs considered 'before and after data' and spatial variation. In this paper, we are interested in assessing the effect of a no-take reserve on the reef fish assemblage in a northwestern Mediterranean example. Data were obtained from a three-year survey using underwater visual censuses (UVC), before and after MPA establishment. Permutational multivariate analysis of variance (PERMANOVA) and multivariate regression trees (MRT) were used to evaluate the effects of reserve protection on the reef fish assemblage, while accounting for habitat. Modelled biological responses were abundances and diversity indices calculated at different levels of the assemblage. Significant effects were found for many of these metrics. In addition to PERMANOVA, univariate models provided more insight into the magnitude and direction of effects. The most sensitive metrics were related to large species and species targeted by fishing. These results may be used to choose the metrics that are more suitable as community-based indicators of MPA impact in the perspective of monitoring programs.

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1. Introduction

Over the last 15 years, most coastal fish resources have been overexploited (Lauck et al., 1998; Castilla, 2000), raising doubts about the long-term sustainability of certain fisheries (Murray et al., 1999; Pauly et al., 2002). In addition, fish habitat has also been strongly altered by widely used fishing

gears such as trawls and dredges, resulting in reduced seabed complexity and removal of macrobenthic organisms that provide shelter for others (Sumaila et al., 2000). Marine protected areas (MPAs) are increasingly considered in coastal areas as an instrument to preserve vagile fauna and habitat from detrimental effects of fishing (Francour et al., 2001; Halpern, 2003; Sainsbury and Sumaila, 2003). The use of

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anti-trawling artificial reefs along the boundaries of several French, Italian and Spanish MPAs has proved to be an effective way of excluding non-selective towed fishing gears which bear detrimental effects on habitats (Harmelin, 2000). It is anticipated that MPAs and, in particular, no-take reserves would be more effective as a fishery and conservation tool for organisms that have relatively sedentary adult life stages and exhibit larval dispersion, enabling biomass exportation to the surrounding areas (Nowlis and Roberts, 1999; Chiappone and Sealey, 2000). There are many documented examples where fished species have benefited from reserve establishment, in particular through increases in mean size and abundance (for reviews, see Roberts and Polunin, 1991; Dugan and Davis, 1993; Rowley, 1994; Bohnsack, 1998; Halpern, 2003). But MPA effects may be diverse in direction and magnitude (Halpern and Warner, 2002). Hence, effects depend on species and timing with respect to reserve establishment (Mosqueira et al., 2000). Sometimes, biological responses (abundance, density, biomass, average size, and diversity of organisms) either consistently increased within the reserve over time (Russ and Alcala, 1996) showed little change over time (Denny and Babcock, 2004), or initially rose but then fell back to original levels (Dufour et al., 1995). Biological responses to protection are also highly variable among fish taxa, but in general species targeted by exploitation are affected in a more positive way than non-target species (Côté et al., 2001), even in the case of recreational fishing (Westera et al., 2003). Large-bodied species also respond more to protection, irrespective of their fishery status (Mosqueira et al., 2000). Most non-target species appear either not to respond to protection (Rakitin and Kramer, 1996) or to respond negatively by showing reduced abundances, perhaps in response to greater predator pressure within reserves (McClanahan et al., 1999).

In spite of these indications, it would be premature to conclude that no-take reserves are always effective for fisheries management, because there are relatively few empirical studies, many of which are poorly designed (Russ, 2002), and even the reported increases in density within reserve borders can be slight (Sale et al., 2005). Attempts to detect explained and predicted effects of MPAs should be based on statistical tests that distinguish between natural variability and the influence of management (Allison et al., 1998; Fraschetti et al., 2002; Benedetti-Cecchi et al., 2003). Despite many empirical studies on reserves, only a few included data collected before and after reserve establishment (Willis et al., 2003), which are the most appropriate for investigating the impact of reserve establishment. Note, however, that if controls are numerous enough to allow an asymmetric comparison with the protected site, this impact can also be detected through “after data” (ACI, After Control Impact) (Glasby, 1997). In addition, few studies account for spatial and temporal variabilities of species, linked to environmental and biological factors other than MPA status (García-Charton and Pérez-Ruzafa, 1999). Fish populations usually exhibit variable degrees of spatial and temporal fluctuation in different parts of the habitat in which they occur. Hence, habitat structure often explains a substantial proportion of the observed variation in fish abundance (García-Charton and Pérez-Ruzafa, 1999; García-Charton et al., 2000; Ferraris

et al., 2005) and if the experimental design fails to capture this variability, any observed differences in fish assemblages may be confounded by differences in habitat (Westera et al., 2003).

An additional problem for studying MPA effects at the fish assemblage level is that most multivariate methods do not test for the presence of interactions and do not measure the magnitude of temporal changes in spatial differences (Clarke, 1993; Underwood and Chapman, 1998). Besides, multivariate analysis of variance requires assumptions about correlations between pairs of variables, assumptions that are rarely met in ecological data sets (Johnson and Field, 1993). Recent developments of permutational multivariate analyses of variance (Anderson, 2001a,b; Anderson and ter Braak, 2003) could help to overcome these problems.

Explicit objectives and monitoring aimed at determining if objectives are met are essential to MPA success (Allison et al., 1998; Claudet and Pelletier, 2004). Monitoring programs provide data for management decisions through the computations of indicators to evaluate progress in conservation programs (Olsen, 2003). Provision of indicators addressing the range of management objectives is needed for integrated coastal management (Belfiore, 2003). These indicators are used for evaluating the effects of interest with respect to management, and for communicating these results to the managers (Linton and Warner, 2003). They must be tailored to particular uses and contexts in both scale and content (Dahl, 2000). Potential ecological indicators can be assessed through their relevance (i.e., the link with the question used) and their effectiveness (which encompasses the issues of precision, accuracy and statistical power) (Nicholson and Fryer, 2002). Many metrics have been used for assessing MPA effects on fish assemblages (for review, see Pelletier et al., 2005), the most frequently used being abundance, biomass, diversity indices and mean size. Pelletier et al. (2005) estimated the relevance and effectiveness of these metrics based on published studies. They showed that in many instances, the use of these metrics led to statistical results that were not significant. For example, the overall species richness of the fish assemblage did not appear to be sensitive to MPA status in the reviewed studies (as also indicated by Russ, 1985; Harmelin et al., 1995). Aside from metrics computed from biological responses encompassing several species or the whole fish assemblage, indicators may be constructed with respect to individual species. The concept of indicator species has been widely used in water management (Bain et al., 2000), but more rarely in marine ecology (Mouillot et al., 2002; Sosa-López et al., 2005). An indicator species is expected to be abundant throughout the studied area and should be easy to sample (Linton and Warner, 2003). Using a large variety of indicator species could provide fine-grained information (Kremen, 1992). To avoid selecting indicator species on expert opinions, intuition and anecdotal information (Saetersdal et al., 2005), Dufrêne and Legendre (1997) developed a flexible and asymmetrical approach based on empirical data to identify indicator species. This method is based on species' specificity and fidelity, combining species' relative abundance and its frequency of occurrence in a group of sites or transects.

In the present study, we are interested in assessing the effects of a no-take marine reserve on the fish assemblage in a Mediterranean ecosystem. Because we were aiming at an evaluation at the assemblage level, while accounting for the assemblage structure, we analysed all data together, and applied multivariate approaches to several metrics pertaining to different groups of fish. To ensure that MPA effects were not confounded with other factors structuring spatial variability of fish, habitat characteristics were considered in the models. Finally, model results were interpreted using a method for identifying indicator species that could be relevant for monitoring and management purposes.

2. Materials and methods

2.1. The studied system

Located in the French northwestern Mediterranean, the Côte Bleue Marine Park (CBMP) was established in 1983. Its primary aim was to protect marine biodiversity, to favour social and economic activities linked to the sea, especially fisheries, and to promote public education and scientific research. The CBMP comprises two effectively enforced no-take reserves: Carry (85 ha), established in 1983, and Couronne (210 ha), established in late 1995 (Fig. 1). In addition to the reserves, two kinds of artificial reefs, for protection against illegal trawling, and for biomass production, were immersed within the park since 1983, several of them being set at the

border of the two reserves to ensure trawl exclusion. In both Carry and Couronne MPAs, fishing, harvesting, scuba diving, anchoring and dredging are forbidden. Compliance is high because the two MPAs were established with the support of users. Commercial and recreational fishing occurs outside the MPAs (Francour et al., 2001). The commercial fishery uses gill nets set on the bottom. They are on average 1.5 m high, and 2500–3500 m long. Gill net is well suited for small in-shore bottoms with chequered patterns of *Posidonia oceanica* meadows, rocks and sand. There are 40 fishing boats and the number of fishers has remained stable over the last 20 years. The recreational fishery comprises 60 sailors on average, mainly uses handline, and targets Labridae and Sparidae. Aside from these fisheries, there is some trawling activity in the vicinity of the CBMP, and occasionally some illegal trawling occurs in the inshore area, explaining immersions of anti-trawling artificial reefs. The aim of this study was to test whether or not the Couronne MPA together with bordering artificial reefs is effective in restoring local fish assemblages.

Fish assemblages of rocky coasts and artificial reefs in the northwestern Mediterranean have been the focus of several studies (Bell and Harmelin-Vivien, 1982; Bell, 1983; Dufour et al., 1995; Harmelin et al., 1995; Harmelin, 1999; Charbonnel et al., 2000, 2002; Jouvenel and Pollard, 2001). Rocky reef fish assemblages are found to be characterised by the dominance of three families, namely the Labridae (*Labrus* and *Ctenolabrus*), the Sparidae (*Diplodus*) and the Serranidae (*Serranus*).

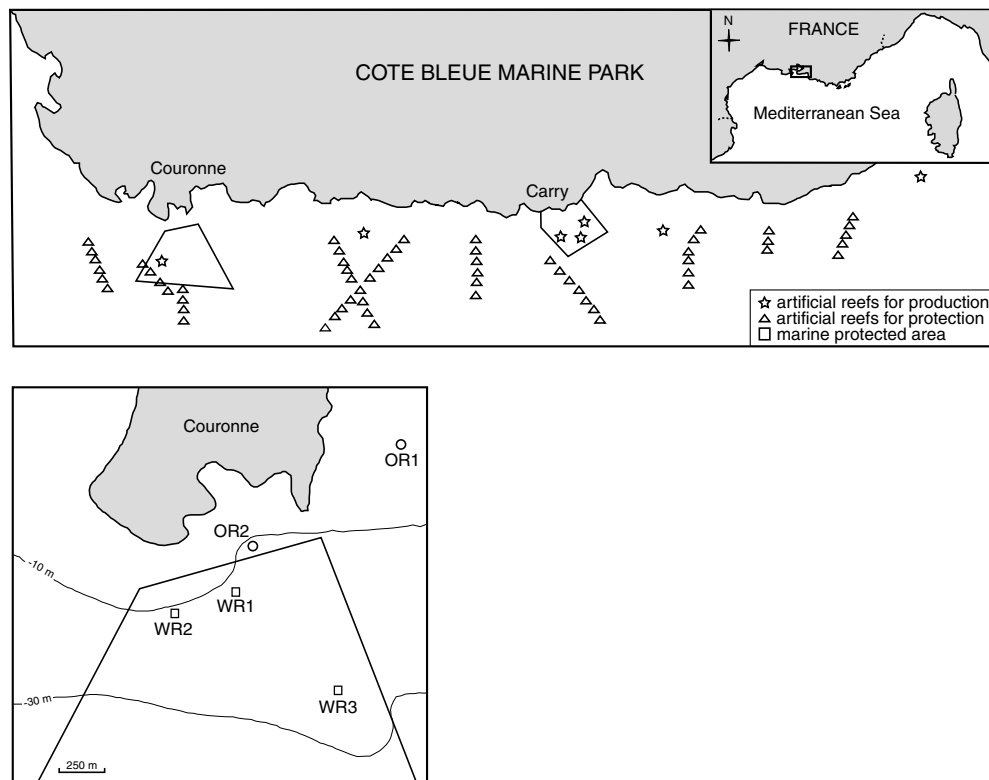


Fig. 1 – The Côte Bleue Marine Park (northwestern Mediterranean) and the Couronne MPA considered in the study. The five surveyed sites were WR1, WR2 and WR3 (within the reserve boundaries) and OR1 and OR2 (outside the reserve).

Many of these species are territorial, with limited home ranges and low inter-annual variations. Sedentary planktivorous fish are also found in the assemblages with Pomacentridae in the shallower areas and Serranidae (*Anthias anthias*) in the deepest ones. Species movements may be in groups but are always restricted in space. They are mainly dictated by seasonal patterns and water temperature. A few pelagic species are also observed in these assemblages during summer, e.g. Carangidae, Clupeidae and some Sparidae species (*Sparus aurata* and *Dentex dentex*).

2.2. Sampling design and data collection

Surveys were conducted at the end of summer during three years: before MPA establishment, in 1995, and after, in 1998 and 2001. Two locations were considered in this study: one within the reserve (WR) boundaries and one outside the reserve (OR). Two sites per location were chosen in the same range of depth (between 14 and 18 m) (Fig. 1). Within the reserve, an additional site was sampled at larger depths (between 24 and 26 m) to study the spatial variation and the effect of depth upon fish abundance.

Underwater visual census (UVC) monitoring techniques provide qualitative and quantitative surveys with a limited impact on the ecosystem, and are therefore particularly suited for marine reserves (Harmelin et al., 1995; Ciriaco et al., 1998). Twelve 20-m-long transects were sampled by scuba divers in each site. Divers swam one way for 8–10 min along each transect, identifying and recording the number and size of each fish species observed within a distance of 2.5 m on each side of the transect. Fish sizes were estimated according to three size groups (small, medium, and large); the total fish abundance of a species being the sum of the abundances per size group. For each species, size groups were defined using 33% and 66% percentiles of the maximum size generally observed in the region. All fish seen were recorded but pelagic species (*Sardina pilchardus*) and notoriously cryptic species (e.g. Gobiidae, Blenniidae, Tripterygiidae) were excluded from the analyses. Sampling methodology and divers remained the same each year to minimise biases inherent in UVC (Kulbicki, 1998; Edgar et al., 2004a). The position of the transects was the same for each year. Under the hypothesis that habitat did not change in the surveyed years, we thus assumed that inter-annual variations in fish assemblage were not caused by habitat changes.

Although the surveyed sites were very similar, environmental data were collected to assess small-scale spatial variability. For each transect, the complexity of the substratum was coded into three classes: 1 for smooth bottoms, 2 for smooth bottoms with a few blocks lower than 50 cm and not suitable for shelter, 3 for bottoms with more blocks, some higher than 1 m, and a lot of refuges. The percent linear cover of *P. oceanica* was estimated along each transect after fish counts. Divers also recorded the depth at the beginning and at the end of each transect.

2.3. Analysis of data

We were interested in assessing whether the Couronne MPA together with bordering artificial reefs is effective at restoring

local fish assemblages. This evaluation was carried out at the fish assemblage level and should thus account for assemblage structure. In order to do so, we used multivariate techniques that are suited for ecological data. In a second step, we ensured that the observed spatial variation was due to MPA effects and not due to other factors structuring spatial variability of fish, by considering habitat characteristics together with MPA design in a single model. Model results were interpreted using a method for identifying indicator species that could be relevant for monitoring and management purposes.

2.3.1. Modelling spatial and temporal variations between within and outside the MPA

Fish abundance was modelled as a function of Location, Year, and Site. All three factors were treated as fixed, the Site factor being nested within the Location factor, Year factor being crossed with the two other factors. For each surveyed year, data were collected at two locations, two sites within each location and $n = 12$ transects within each site, leading to a total of 144 observations in the data set. Because depth strongly structures the spatial distribution of fish, data from the deeper site WR3 within the MPA were not considered in the present model (but see Section 2.3.2), in order to maximise the probability of detecting an impact of MPA (Benedetti-Cecchi, 2001). To estimate synergisms and antagonisms among the effects of the different factors investigated (Underwood, 1981), the model included all combinations of the factor levels. Given the design, the linear algebraic model thus wrote:

$$X_{ijkz} = \mu + Y e_i + L o_j + Y e \times L o_{ij} + S i(L o)_{k(j)} + Y e \times S i(L o)_{ik(j)} + e_{z(ijk)},$$

where X_{ijkz} represents the set of abundances observed at the z th replicate ($z = 1, 2, \dots, 12$) of the k th level of the nested factor Site (Si) in the j th level of the factor Location (Lo) crossed with the i th level of the factor Year (Ye). μ represents the overall mean abundance vector. $Y e_i$ represents the effect of the i th level of the factor Year ($i = 1, 2, 3$); $L o_j$ denotes the effect of the j th level of the factor Location ($j = 1, 2$); $S i(L o)_{k(j)}$ represents the effect of the k th level of the nested factor Site in the j th level of the factor Location ($k = 1, 2$); $Y e \times L o_{ij}$ and $Y e \times S i(L o)_{ik(j)}$ correspond to the interaction effects of the factors Year and Location and of the factors Year and Site. Finally, $e_{z(ijk)}$ represents the error term associated with each observation. All factors being fixed, the term used for denominator mean square in the F-ratio was thus always the residuals for all terms in the model.

With Before–After–Control–Impact data, the consequences of the MPA on the fish assemblage may be studied from differences in the change of fish species biological responses (e.g., abundance, richness, and diversity) inside the MPA location from before to after its establishment compared with such changes from before to after in the control location (sensu Underwood, 1993). Under this model, an effect of the reserve on the fish assemblage is evidenced if the interaction term between Year and Location is statistically significant. Such differences in biological responses across years and between inside and outside the MPA (termed inside/outside differences from this point onwards)

were interpreted by pair-wise comparisons conducted on these interaction terms. Differences between sites over years in a given location (i.e., a significant Year \times Site(Location) interaction) do not interfere with the MPA effects. These differences could be due to small-scale variability in the assemblages of fish. The presence of significant higher-order interactions warns that the experimental treatments do not operate independently, in any combinations (Underwood, 1981).

The observed fish assemblage comprised 40 species (Appendix 1). In order to analyse how the fish assemblage responds to the MPA-artificial reef system, models were constructed for sets of abundance indices calculated at several levels and for several components of the fish assemblage (see Appendix 1): (i) abundance per species for the whole fish assemblage; (ii) abundance per observed size group (small, medium and large); (iii) abundance per species for species grouped by species size; (iv) abundance per species for species groups based on the species' fishing value; and (v) abundance per species for species groups based on mobility. Species size groups were based on the minimum and maximum lengths generally observed for each species in the northwestern Mediterranean, i.e. 8–20 cm, 20–30 cm and 30–200 cm. For fishing value, three groups of species were considered: unfished species, species with low fishing value, and species with medium to high fishing value. Regarding mobility, we distinguished mobile species and sedentary species; mobile species including demersal species displaying horizontal movements and possibly vertical movements, and sedentary species including benthic species and species that move only marginally, both horizontally and vertically.

Unfortunately, distributions of abundances per fish species are usually highly skewed and contain many zeros. Conventional multivariate inferential methods such as MANOVA are not appropriate for this kind of data. We thus used the permutational multivariate analysis of variance (PERMANOVA), initially called NPMANOVA (Anderson, 2001a; McArdle and Anderson, 2001). This method analyses the variance of multivariate data explained by a set of explanatory factors on the basis of any distance or dissimilarity measure of choice, thereby allowing for a wide range of empirical data distributions. The method provides *P*-values by permutations, so that effects linked to each factor or interaction between factors may be tested in a more robust way than with MANOVA. To perform the PERMANOVA, the FORTRAN computer program DISTLM4 was used (Anderson, 2004b). In the models, fish abundance data were log-transformed. We used the binomial deviance dissimilarity that is appropriate for this kind of empirical distributions (Anderson and Millar, 2004). Each term in the model was tested through permutation tests based on 4999 permutations of residuals under a reduced model to obtain *P*-values. This permutation method is generally thought to be best suited because it provides the best statistical power and the most accurate Type I error (Anderson and Legendre, 1999). When significant at the 0.05 level, the Ye \times Lo_{ij} interaction term was investigated through a posteriori pair-wise comparisons using 4999 random permutations to obtain *P*-values. Because multiple multivariate interactions are difficult to visualise, they were projected in the two-dimensional plane generated by the first

two axes obtained from discriminant analyses conducted separately for each year to discriminate fish abundances observed within the reserve from those observed outside the reserve. This way, within-location variabilities can be visually compared and tested. In addition, the correlation between the fish species and the axis indicates which species are determined in explaining the differences between locations, i.e., which species exhibit spatial differences in abundance due to MPA.

In practice, discriminant analyses were achieved using the CAP software (Anderson, 2004a) which calculates a canonical analysis on the principal coordinates based on any symmetric distance matrix, including a permutation test (Anderson, 2004a). We used the same data transformation and dissimilarity measure for CAP and for PERMANOVA, i.e., abundance data were log-transformed, the distance measure used was the binomial deviance dissimilarity and tests relied on 4999 permutations.

Previous multivariate analyses yield a test of the MPA effect and allow to identify species that are closely linked to these effects. Furthermore, we were also interested in analysing the effect of the MPA on diversity metrics. This was addressed through univariate analyses. We first modelled the abundance of two fished species with high fishing value and one species with low fished value. We selected species encountered in more than 50% of the transects across the factors of interests (see Appendix 1), namely *Coris julis*, *Serranus cabrilla*, and *Symphodus doderleini*. We then modelled the overall fish abundance, species richness and the Shannon-Wiener diversity index. For each of these variables, two analyses were carried out: one considering all fish and the other considering only large fish, because large fish usually respond more to protection (Mosqueira et al., 2000). Analyses were conducted using permutation tests realised using the DISTLM4 software (Anderson, 2004b) with 4999 random permutations. In the models, only abundance variables were log-transformed. Unlike multivariate analyses described above, we used a Euclidean distance in the univariate models. We particularly tested the Ye \times Lo_{ij} interaction term from a posteriori pair-wise comparisons, based on 4999 random permutations under a 0.05 significance level. Boxplots were used to illustrate mean abundances per Location and Year for each modelled variable.

2.3.2. Incorporating the influence of depth and habitat in the assessment of MPA effects

The models described in the previous subsection do not account for environmental variables such as habitat and depth, measured as part of the study design. Depth issues were avoided by excluding a site that was deeper than the others.

A first appraisal of the relationships between species abundance and environment is provided by modelling the abundances of all species as a function of three environmental covariables: mean depth, percentage linear cover of *P. oceanica*, and substrate complexity. This was achieved by a multivariate analysis of covariance. This analysis was carried out from the DISTLM4 software. In the model, the distance measure was the binomial deviance dissimilarity and 4999 permutations were done for the tests.

In a second step, only significant environmental covariables and factors allowing the evaluation of MPA effects on the set of abundances per species were included as explicative variables of fish abundance data per species and per transect. In this purpose, we used multivariate regression trees (MRT) (De'ath, 2002). This multivariate discrimination technique constructs a hierarchical tree through successive dichotomies of the set of observations. It was used to build hierarchical groups of observations to analyse which of the protection or the environment affected more the fish assemblages. Splits and clusters are characterised by values and conditions on explicative variables. Fish abundance data were log-transformed. The MRT technique does not require any assumptions about the form of the relationships between observations and explicative variables. Trees were pruned by cross-validation using the minimum rule of Breiman et al. (1984). These analyses were done using the mvpart package of the R statistical software (Therneau et al., 2004). Two MRT were calculated from log-transformed abundance data concerning: (i) the four sites previously used in the PERMANOVA (WR1, WR2, OR2 and OR2), and (ii) the four previous sites and in addition the deeper site within the MPA, WR3 (Fig. 1), where the resulting number of sampling units was 180. The site WR3 was included to analyse the respective influence of depth and protection upon fish assemblages. In each case (with and without WR3) two MRT

were calculated, one with the fish abundance data of all sizes and one only with large fish. Explanatory variables were the three factors Year, Location, Site, and the three environmental variables (percentage linear coverage of *P. oceanica*, substrate complexity and average depth). Two fish species that display schooling behaviour were excluded from MRT calculations (*Boops boops* and *Chromis chromis*).

2.3.3. Identifying indicator species

In a last step, the MRT were used to identify indicator species. Tree leaves, i.e., clusters corresponding to a given split, were characterised by species using the indicator value (IndVal) method (Dufrene and Legendre, 1997). With this approach, indicator species characterise a cluster of observations corresponding to a given leaf of the tree if it is simultaneously abundant and frequent in the group compared to the whole set of observations. The index used to identify indicator species is the product of relative abundance and relative frequency of occurrence. It is maximum for a given cluster when the species is found in all observations in this cluster and is not encountered in other clusters. The statistical significance of a species as an indicator at the 0.05 level was evaluated using a randomisation procedure. Calculations were done using the IndVal 2.0 FORTRAN computer program.

Table 1 – PERMANOVA table of abundances per fish species conducted on fish size group

Fish considered	Source of variation	df	SS	F	P
All (40 variables)	Year	2	45.29	5.65	0.0002***
	Location	1	47.37	11.82	0.0002***
	Site(Location)	2	35.68	4.45	0.0002***
	Ye × Lo	2	56.06	6.99	0.0002***
	Ye × Si(Lo)	4	33.15	2.07	0.0130*
	Residual	132	528.95		
Large (40 variables)	Ye	2	21.69	6.04	0.0004***
	Lo	1	2.54	17.02	0.0002***
	Si(Lo)	2	7.54	2.10	0.0718, n.s.
	Ye × Lo	2	1.88	5.82	0.0004***
	Ye × Si(Lo)	4	8.34	1.16	0.3272, n.s.
	Residual	132	236.81		
Medium (40 variables)	Ye	2	25.39	4.66	0.0012**
	Lo	1	19.36	7.10	0.0002***
	Si(Lo)	2	21.56	3.96	0.0030**
	Ye × Lo	2	29.39	5.39	0.0002***
	Ye × Si(Lo)	4	16.78	1.54	0.1316, n.s.
	Residual	132	359.74		
Small (40 variables)	Ye	2	5.92	3.72	0.0038**
	Lo	1	1.06	1.33	0.2886, n.s.
	Si(Lo)	2	5.61	3.52	0.0052**
	Ye × Lo	2	4.21	2.65	0.0256*
	Ye × Si(Lo)	4	6.21	1.95	0.0438*
	Residual	132	104.99		

n.s., not significant.

PERMANOVAs were based on the binomial deviance dissimilarity measure. P-values were obtained using 4999 permutations of residuals under a reduced model.

* $P < 0.05$.

** $P < 0.01$.

*** $P < 0.001$.

3. Results

3.1. Modelling spatial and temporal variations within and outside the MPA

PERMANOVA of the fish abundances per species showed a significant multivariate interaction between the factors Year and Location whatever the fish size was (Table 1); thereby indicating an MPA effect. There were no initial differences in abundances between locations (within vs. outside reserve) before MPA establishment, in 1995 (Table 2). Application of the discriminant analysis to abundance of large fish showed that the classification in two groups was not significant in 1995 ($P = 0.5456$; the proportion of classification in the correct cluster was only 58%), confirming the absence of inside/outside differences in the assemblages of large fish before MPA establishment.

To the exception of small fish individuals, inside/outside differences in species abundance became significant after MPA establishment, abundances being on average higher within the reserve (Appendix 2). These differences were more significant for large fish than for medium-sized fish. The results of discriminant analyses of the abundance of large fish illustrated that these differences appeared after 1995 (Fig. 2). A similar, although less marked, trend could be observed for medium-sized fish and for all fish (not reported here). After MPA establishment, i.e., in 1998 and 2001, the classification of fish abundance data into two groups became significant ($P = 0.0002$ in both cases). For both years, 83% of observations of the original 48 were correctly classified. In 1998 (results not reported in Fig. 2), only species positively correlated with the canonical axes showed high correlations (>0.5), namely large fish of the commercial species *S. cabrilla*, *C. julis*, *Ctenolabrus rupestris*, *S. doderleini* and

Table 2 – P-values for pair-wise comparisons conducted after PERMANOVAs of fish species abundance data

Variables	1995: WR vs. OR, P	1998: WR vs. OR, P	2001: WR vs. OR, P
All fish	0.0896, n.s.	0.0002***	0.0002***
Large fish	0.4688, n.s.	0.0002***	0.0002***
Medium fish	0.2010, n.s.	0.0008***	0.0004***
Small fish	0.1176, n.s.	0.1910, n.s.	0.0786, n.s.
Large species (30–200 cm)	0.9998, n.s.	0.1776, n.s.	0.0046**
Medium species (20–30 cm)	0.9164, n.s.	0.0002***	0.0004***
Small species (8–20 cm)	0.0376*	0.0198*	0.0016**
Low value commercial species	0.0182*	0.0002***	0.0004***
Medium to high value commercial species	0.6396, n.s.	0.0032**	0.0002***
Mobile species	0.8100, n.s.	0.0004***	0.0004***
Sedentary species	0.0200*	0.0002***	0.0002***

n.s., not significant.

Comparisons were performed for inside/outside differences for each year.

Only the metrics for which the interaction (Year \times Location) was significant are reported (see Tables 1, 3, 4, and 5). WR: within reserve; OR: outside reserve. P-values were obtained using 4999 permutations. The pair-wise tests have not been corrected for multiple comparisons.

* $P < 0.05$.

** $P < 0.01$.

*** $P < 0.001$.

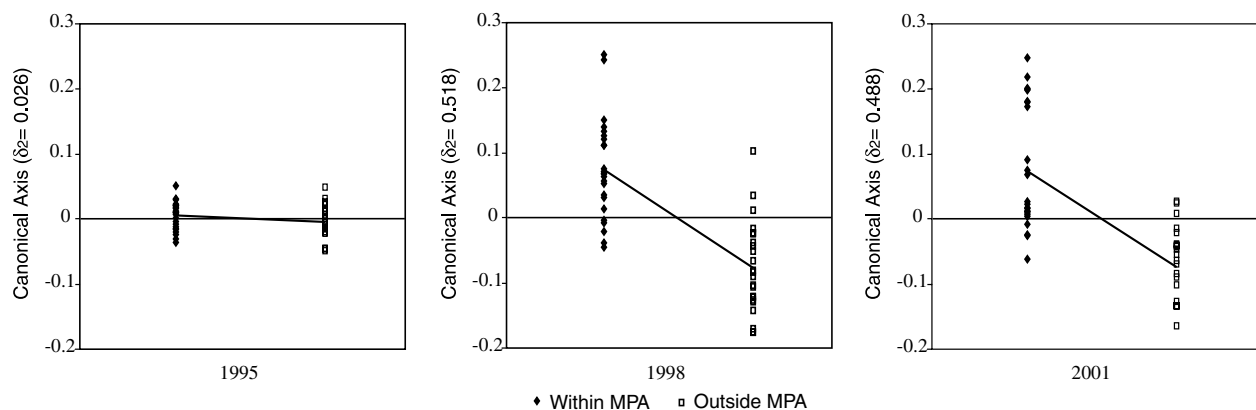


Fig. 2 – Discriminant analyses plots for each year based on the binomial deviance dissimilarity measure. Analyses compared the fish assemblages in the two locations: within the MPA boundaries (black diamonds) and outside the MPA (white rectangles). There are $n = 24$ observations in each location for each year.

Symphodus mediterraneus. In 2001, high positive correlations with the canonical axes were observed for large *Diplodus sargus*, *Diplodus vulgaris*, *Symphodus melanocercus*, *Symphodus tinca*, and again *S. cabrilla*, *C. julis*, *C. rupestris* and *S. doderleini*; all being commercial species. In both cases, large positive correlations indicate significantly larger abundances for these species within the MPA. However, negative correlations indicating higher abundances outside the reserve were observed for *Spicara maena* and *C. chromis*. Between sites variability was only significant for small fish outside the MPA in 2001 (pair-wise comparison, $P = 0.0124$).

In the case of species size groups, PERMANOVAs showed a significant Year \times Location interaction for all groups (Table 3). Inside/outside differences were significant across all years only for small species (Table 2), abundances being on average always higher within the MPA (Appendix 2). For medium-sized species, differences between locations became significant after MPA establishment, and only in 2001 for large species (see Appendix 2 for mean abundances). Between-sites differences could be evidenced for large and medium-sized fish species. These small-scale differences were significant within the reserve in 2001 for large species (pair-wise comparison, $P = 0.0010$) and within the reserve in 1998 and outside the MPA in 2001 for medium-sized species (pair-wise comparison, $P = 0.0320$ and 0.0010 , respectively).

There was no significant Year \times Location interaction for unfished species (Table 4). The abundances of unfished species were different between years (Table 4) and only significantly different between 1998 and 2001 (pair-wise comparison, $P = 0.0196$). For fished species, PERMANOVAs produced a significant Year \times Location interaction. The abundances of species with low fishing value were significantly

different between locations across all years. Significant inside/outside differences occurred only after MPA establishment for species of medium to high fishing value and became more significant over the years (Table 2). Across all years, the sites inside the reserve showed on average lower abundances of species with low fishing value (Appendix 2). On the opposite, species with medium to high fishing value were, on average, more abundant inside the reserve. For these species, there were significant between-sites difference outside the MPA in 2001 (pair-wise comparisons, $P = 0.0018$).

PERMANOVAs conducted on the abundance per species when species were grouped by mobility showed a significant interaction for both mobile and sedentary species (Table 5). Inside/outside differences were significant across all years, for these species, except in 1995 where there were no differences for mobile species (Table 2). For both groups of species, when inside/outside differences were significant, abundances were on average higher inside the MPA (Appendix 2). Significant between-sites differences occurred only for sedentary species in 1998 within the MPA (pair-wise comparison, $P = 0.0140$) and in 2001 outside the MPA (pair-wise comparison, $P = 0.0060$).

Permutational univariate ANOVAs on overall abundance, richness and diversity showed a significant Year \times Location interaction for all metrics considered (Table 6). Inside/outside differences were not significant before MPA establishment (Table 7). In 1998, the differences between locations on overall abundance, richness and diversity became significant only for large fish. In 2001, inside/outside differences in overall abundance, species richness and diversity were all significant whether considering all or large fish. When the differences were significant, the value of the metric was always larger within the MPA (Fig. 3), except for the diversity

Table 3 – PERMANOVA table of abundance per species when analyses were conducted per species size group

Species sizes	Source of variation	df	SS	F	P
Large 30–200 cm (15 species)	Year	2	0.10	0.55	0.6894, n.s.
	Location	1	0.06	0.62	0.5640, n.s.
	Site(Location)	2	0.94	5.00	0.0024**
	Ye \times Lo	2	1.04	5.52	0.0002***
	Ye \times Si(Lo)	4	1.42	3.79	0.0008***
	Residual	132	12.38		
Medium 20–30 cm (10 species)	Ye	2	1.13	0.71	0.5920, n.s.
	Lo	1	14.23	17.88	0.0002***
	Si(Lo)	2	16.10	10.11	0.0002***
	Ye \times Lo	2	13.70	8.61	0.0002***
	Ye \times Si(Lo)	4	10.81	3.40	0.0020**
	Residual	132	105.10		
Small 8–20 cm (15 species)	Ye	2	1.19	9.91	0.0002***
	Lo	1	8.18	8.03	0.0008***
	Si(Lo)	2	1.94	0.95	0.4592, n.s.
	Ye \times Lo	2	10.24	5.02	0.0012**
	Ye \times Si(Lo)	4	7.36	1.81	0.0986, n.s.
	Residual	132	134.52		

n.s., not significant.

PERMANOVAs were based on the binomial deviance dissimilarity measure. P-values were obtained using 4999 permutations of residuals under a reduced model. The range of species sizes is reported for each group.

* $P < 0.05$.

** $P < 0.01$.

*** $P < 0.001$.

Table 4 – PERMANOVA table of abundance per species when analyses were conducted per species group based on fishing value

Fishing value	Source of variation	df	SS	F	P
Unfished (5 species)	Year	2	2.89	3.95	0.0252*
	Location	1	0.75	2.05	0.1824, n.s.
	Site(Location)	2	0.62	0.85	0.4704, n.s.
	Ye × Lo	2	1.50	2.05	0.1496, n.s.
	Ye × Si(Lo)	4	2.89	1.98	0.1174, n.s.
	Residual	132	48.20		
Low (13 species)	Ye	2	13.45	8.55	0.0002***
	Lo	1	8.93	11.36	0.0002***
	Si(Lo)	2	8.79	5.59	0.0002***
	Ye × Lo	2	7.44	4.73	0.0006***
	Ye × Si(Lo)	4	3.64	1.16	0.3462, n.s.
	Residual	132	103.83		
Medium to high (22 species)	Ye	2	1.42	1.30	0.3008, n.s.
	Lo	1	9.95	18.18	0.0002***
	Si(Lo)	2	4.99	4.56	0.0028**
	Ye × Lo	2	11.72	10.71	0.0002***
	Ye × Si(Lo)	4	6.22	2.84	0.0086**
	Residual	132	72.26		

n.s., not significant.

PERMANOVAs were based on the binomial deviance dissimilarity measure. P-values were obtained using 4999 permutations of residuals under a reduced model.

* $P < 0.05$.** $P < 0.01$.*** $P < 0.001$.**Table 5 – PERMANOVA table of abundance per species when analyses were conducted per species group based on mobility**

Species mobility	Source of variation	df	SS	F	P
Mobile species (18 species)	Year	2	7.99	3.02	0.0214*
	Location	1	11.23	8.49	0.0002***
	Site(Location)	2	11.69	4.42	0.0018**
	Ye × Lo	2	15.75	5.95	0.0002***
	Ye × Si(Lo)	4	7.46	1.41	0.2276, n.s.
	Residual	132	174.55		
Sedentary species (22 species)	Ye	2	16.54	8.68	0.0002***
	Lo	1	15.19	15.93	0.0002***
	Si(Lo)	2	7.34	3.85	0.0014**
	Ye × Lo	2	14.69	7.70	0.0002***
	Ye × Si(Lo)	4	8.37	2.19	0.0158*
	Residual	132	125.85		

n.s., not significant.

PERMANOVAs were based on the binomial deviance dissimilarity measure. P-values were obtained using 4999 permutations of residuals under a reduced model.

* $P < 0.05$.** $P < 0.01$.*** $P < 0.001$.

index which was higher outside the MPA in 1998 (Fig. 3). However, diversity became lower outside the MPA in 2001. It is interesting to note that for large fish, inside/outside differences were significant from 1998 and increased in 2001. In 1995, total abundance and diversity displayed significant between-sites differences, respectively, inside the MPA (pair-wise comparison, $P = 0.0434$) and outside the MPA (pair-wise comparison, $P = 0.0078$). In 2001, between-sites differences were significant outside the MPA for total abundance, spe-

cies richness, overall diversity and diversity of observed large fish (pair-wise comparison, respectively, $P = 0.0158$, 0.0048, 0.0004, 0.0012).

The permutational univariate ANOVAs of the abundance of two species of high fishing value, *C. julis* and *S. cabrilla*, and one species with low fishing value, *S. doderleini*, revealed a significant Year × Location interaction for the three species (Table 8). The interaction was significant for total species abundance and for the abundance of large fish. In all cases,

Table 6 – ANOVA table for permutational univariate analyses of total abundance, species richness and diversity metrics

Community metrics	Source of variation	df	SS	F	P
<i>Total fish abundance</i>					
All fish	Year	2	6.61	3.08	0.0496*
	Location	1	12.57	11.69	0.0008***
	Site(Location)	2	6.25	2.91	0.0564, n.s.
	Ye × Lo	2	14.15	6.58	0.0024**
	Ye × Si(Lo)	4	27.74	6.45	0.0006***
	Residual	132	141.94		
Large fish	Ye	2	2.36	0.99	0.3838, n.s.
	Lo	1	26.21	21.94	0.0002***
	Si(Lo)	2	7.12	2.99	0.0530, n.s.
	Ye × Lo	2	10.51	4.41	0.0134*
	Ye × Si(Lo)	4	6.41	1.34	0.2602, n.s.
	Residual	132	157.68		
<i>Species richness</i>					
All fish	Ye	2	92.62	6.38	0.0030**
	Lo	1	51.36	7.08	0.0092**
	Si(Lo)	2	27.22	1.88	0.1530, n.s.
	Ye × Lo	2	236.93	16.33	0.0002***
	Ye × Si(Lo)	4	106.11	3.66	0.0064**
	Residual	132	957.50		
Large fish	Ye	2	32.76	2.87	0.0624, n.s.
	Lo	1	156.25	27.34	0.0002***
	Si(Lo)	2	10.00	0.88	0.4190, n.s.
	Ye × Lo	2	140.79	12.32	0.0002***
	Ye × Si(Lo)	4	34.33	1.50	0.2008, n.s.
	Residual	132	754.50		
<i>Diversity</i>					
All fish	Ye	2	1.83	2.74	0.0674, n.s.
	Lo	1	0.82	2.45	0.1332, n.s.
	Si(Lo)	2	8.97	13.41	0.0002***
	Ye × Lo	2	6.92	10.35	0.0004***
	Ye × Si(Lo)	4	5.36	4.01	0.0052**
	Residual	132	44.13		
Large fish	Ye	2	4.80	4.44	0.0136*
	Lo	1	10.17	18.80	0.0002***
	Si(Lo)	2	1.15	1.07	0.3498, n.s.
	Ye × Lo	2	10.78	9.96	0.0002***
	Ye × Si(Lo)	4	5.91	2.73	0.0326*
	Residual	132	71.40		

n.s., not significant.

Permutational univariate ANOVAs were based on Euclidean distances. P-values were obtained using 4999 permutations of residuals under a reduced model.

* $P < 0.05$.

** $P < 0.01$.

*** $P < 0.001$.

inside/outside differences were not significant before MPA establishment (Table 7). In 1998, all inside/outside differences were significant, except for total abundance of *C. julis*. Note that the inside/outside difference was less marked for *S. doderleini*. In 2001, inside/outside differences were still significant for *S. cabrilla* and *C. julis* (and indeed more significant than in 1998 for total abundance of *C. julis*), whereas they were not any longer significant for *S. doderleini*. These differences corresponded to higher abundances inside the MPA compared to outside the MPA (Fig. 4). The results were similar for all the observed *Symphodus* species (eight species), for both total and large fish abundance. Note that between-sites differences

were significant for *C. julis* outside the MPA in 2001 (pair-wise comparison, $P = 0.0026$).

3.2. Joint effects of MPA, depth and habitat on the fish assemblages

A multivariate non-parametric analysis of covariance was performed on the whole fish assemblage to explore the relationship between fish abundance and a set of the three environmental variables, i.e., mean depth, complexity, and linear percentage cover of *P. oceanica*. These covariables were found to have a significant effect on fish abundance ($P = 0.002$). How-

Table 7 – P-values for pair-wise comparisons conducted after permutational univariate ANOVAs of total abundance, species richness and diversity metrics and abundance of two fished species (*Coris julis* and *Serranus cabrilla*) and one species with low fishing value (*Symphodus doderleini*)

Variables	1995: WR vs. OR, P	1998: WR vs. OR, P	2001: WR vs. OR, P
Number of fish	0.7110, n.s.	0.2744, n.s.	0.0038**
Number of fish (large fish individuals)	0.7740, n.s.	0.0004***	0.0002***
Species richness	0.1344, n.s.	1.0000, n.s.	0.0002***
Species richness (large fish individuals)	0.3702, n.s.	0.0006***	0.0002***
Diversity	0.3536, n.s.	0.3790, n.s.	0.0008***
Diversity (large fish individuals)	0.3796, n.s.	0.0012**	0.0002***
<i>Coris julis</i> (all fish)	0.9326, n.s.	0.1256, n.s.	0.0002***
<i>Coris julis</i> (large fish individuals)	0.2550, n.s.	0.0002***	0.0002***
<i>Serranus cabrilla</i> (all fish)	0.9480, n.s.	0.0002***	0.0028**
<i>Serranus cabrilla</i> (large fish individuals)	0.0736, n.s.	0.0002***	0.0002***
<i>Symphodus doderleini</i> (all fish)	0.2160, n.s.	0.0024**	0.1394, n.s.
<i>Symphodus doderleini</i> (large fish individuals)	0.4532, n.s.	0.0022**	0.6334, n.s.

n.s., not significant.
 Comparisons were performed for inside/outside differences for each year.
 Only the metrics for which the interaction (Year \times Location) was significant are reported (see Tables 6 and 8). WR: within reserve; OR: outside reserve. P-values were obtained using 4999 permutations. The pair-wise tests have not been corrected for multiple comparisons.
 * P < 0.05.
 ** P < 0.01.
 *** P < 0.001.

ever, accounting for these covariables including Year, Site and Location factors in the model did not change the significance of the interactions Year \times Location and Year \times Site(Location); respectively, P = 0.002 and 0.0118.

MRT were calculated for abundance per species, in the case of total abundance and abundance of large fish, using previous environmental variables (depth, complexity, and linear percentage cover of *P. oceanica*), and factors Year, Location, and Site, as explanatory variables. In both cases, MRT were calculated considering or not data collected in the deeper site WR3. Although the results were valid in the four cases (total/large fish abundance combined with/without R3), they were only reported for abundance of large fish considering WR3, for the sake of concision (Fig. 5). The first split separated observations within the MPA from observations without the MPA in three cases out of four, the latter corresponding to total abundance with data from site R3. In all three cases, the second split, whether within or outside the MPA, separated 1995 from 1998 and 2001 (i.e., the year before the MPA establishment from the years after). To analyse whether the first split of these trees was indicating a habitat effect or a reserve effect, additional trees were computed using a composite factor for the interaction between factors Year and Location (six levels, e.g. WR.1995). In this case, the first split separated observations within the MPA after its establishment from observations of both locations before MPA establishment and observations from outside the MPA. We thus concluded that the first split indicated a reserve effect. All following splits were similar to the ones obtained in other trees. In the fourth case, i.e., for total abundance with data from site WR3, the first split separated this site from all the others. In the second split, sites were separated depending on whether they were within or outside the MPA. Later splits were similar to the three other cases. Transects within WR3 were distinguished by *P. oceanica* cover (with a percent level of discrimination of 42.5). In the

MRT obtained for the abundance of large fish considering data from site WR3 (Fig. 5), abundances after MPA establishment (1998 and 2001) within MPA were distinguished according to the depth. At shallower depths (<21.5 m), abundances differed between 1998 and 2001. At larger depths (≥ 21.5 m), *P. oceanica* cover explained differences between abundances obtained at distinct transects. Thus, habitat variables were only discriminant after MPA establishment and within the MPA, and differences in abundance over years were more marked in shallower depths than in larger depths.

3.3. Indicator species

Indicator species were searched for the MRT reported in Fig. 5. Two groups of species were indicators of the first split, i.e., separating within from outside the MPA locations, irrespective of other factors: *Apogon imberbis* was specific of observations outside the MPA (i.e., being more abundant and more frequent in these transects, group III in Fig. 5). The Serranidae *S. cabrilla* and the Labridae *C. julis* and *C. rupestris* were specific of observations inside the MPA (group II in Fig. 5). These three species had still significant indicator values after the MPA establishment (i.e., after other splits) but these values were maximum for group II. Outside the MPA, the Labridae *S. doderleini* and *Symphodus roissali* were indicator species of the year before MPA establishment (group F in Fig. 5). Within the MPA, the Sparidae *Sarpa salpa* and the Labridae *S. tinca* were indicator species for years after MPA establishment, whatever the habitat characteristics were (group IV in Fig. 5). No indicator species was found for shallower depths; whereas at larger depths, the Scorpaenidae *Scorpaena porcus* and the Mullidae *Mullus surmuletus* were significant indicator species (group V in Fig. 5). The Serranidae *A. anthias* and the Sparidae *D. vulgaris* and *D. sargus* were specific of MPA, after its establishment, but only at

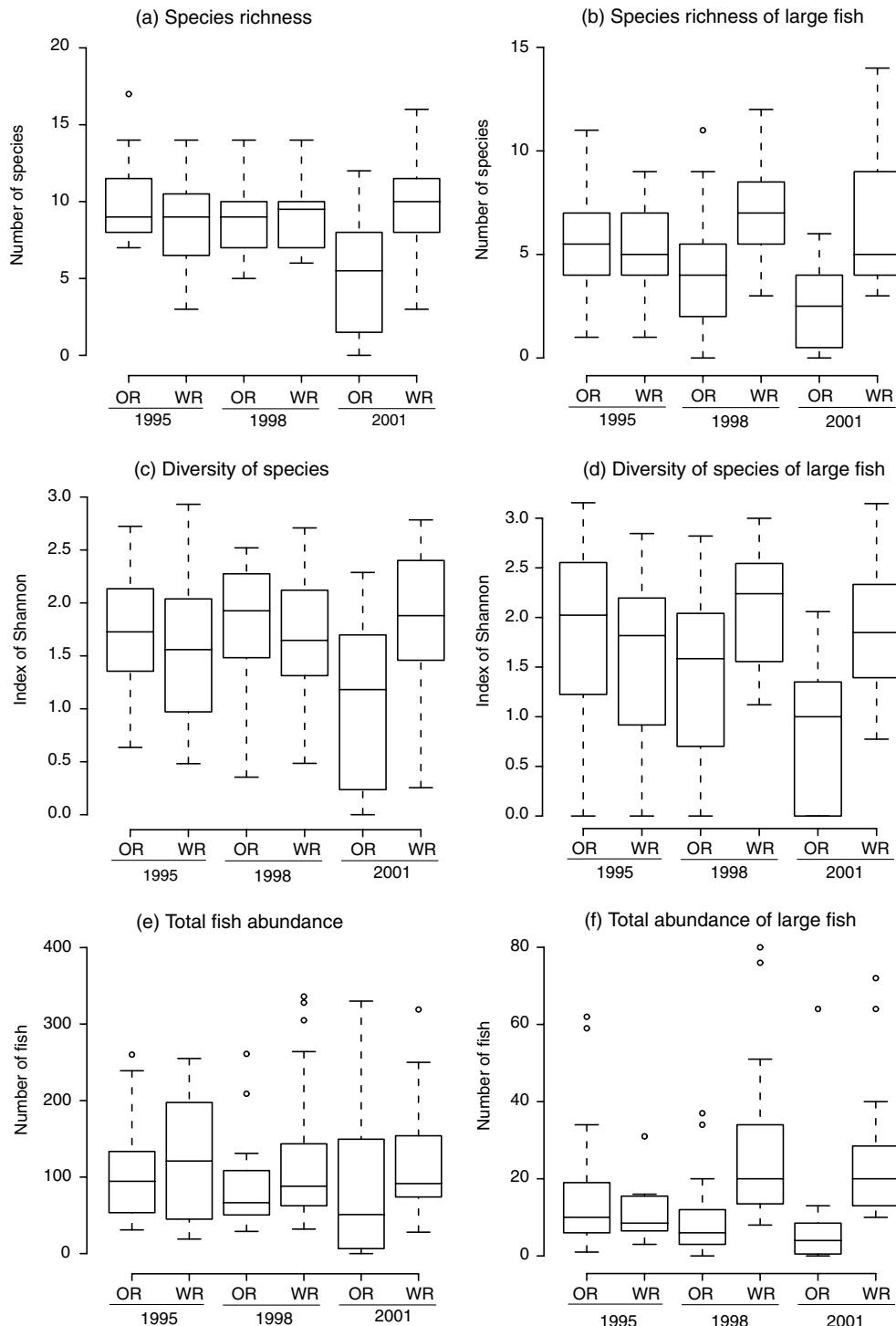


Fig. 3 – Boxplots of (a) species richness, (b) species richness of large fish, (c) diversity index of Shannon, (d) diversity index of Shannon of large fish, (e) overall fish abundance, and (f) overall abundance of large fish for each combination of the levels of factors Year and Location. There were $n = 24$ transects per combination. For better readability, outlying values were not reported on (e) and (f): for total fish abundance, WR.1995: 534, OR.1998: 553; WR.2001: 852 and 3095 fishes; for total abundance of large fish: OR.1995: 105, and 217; WR.1995: 113, 208, 215 and 511 fishes; OR.1998: 243; WR.1998: 158; OR.2001: 106 and 313 fishes.

larger depths and where the percent cover of *P. oceanica* was low (group B in Fig. 5). Species of group A were distinguished from those in group B only by a higher percentage linear

cover of *P. oceanica*; these were namely: the Scorpaenidae *Scorpaena scrofa* and *Scorpaena notata*, the Centracanthidae *S. maena* and the Labridae *Labrus merula* and *Labrus viridis*.

Table 8 – ANOVA table for permutational univariate analyses of the abundance of two fished species (*Coris julis* and *Serranus cabrilla*) and one species with low fishing value (*Symphodus doderleini*)

Fish species	Source of variation	df	SS	F	P
<i>Coris julis</i>					
All fish	Year	2	0.22	0.40	0.6730, n.s.
	Location	1	8.37	29.86	0.0002***
	Site(Location)	2	5.95	10.62	0.0002***
	Ye × Lo	2	12.02	21.46	0.0002***
	Ye × Si(Lo)	4	11.62	10.37	0.0002***
	Residual	132	36.99		
Large fish	Ye	2	4.56	10.47	0.0004***
	Lo	1	24.14	110.77	0.0002***
	Si(Lo)	2	4.96	11.39	0.0002***
	Ye × Lo	2	8.60	19.74	0.0002***
	Ye × Si(Lo)	4	1.78	2.04	0.0880, n.s.
	Residual	132	28.76		
<i>Serranus cabrilla</i>					
All fish	Ye	2	1.09	2.73	0.0710, n.s.
	Lo	1	6.02	2.06	0.0002***
	Si(Lo)	2	4.74	11.85	0.0004***
	Ye × Lo	2	3.33	8.32	0.0002***
	Ye × Si(Lo)	4	1.15	1.44	0.2364, n.s.
	Residual	132	26.43		
Large fish	Ye	2	3.13	10.29	0.0004***
	Lo	1	9.00	59.20	0.0002***
	Si(Lo)	2	0.70	2.29	0.1058, n.s.
	Ye × Lo	2	9.40	2.92	0.0002***
	Ye × Si(Lo)	4	0.58	0.95	0.4312, n.s.
	Residual	132	1.06		
<i>Symphodus doderleini</i>					
All fish	Ye	2	2.08	3.74	0.0252*
	Lo	1	1.49	5.38	0.0238*
	Si(Lo)	2	1.09	1.96	0.1536, n.s.
	Ye × Lo	2	2.72	4.90	0.0104*
	Ye × Si(Lo)	4	1.28	1.15	0.3348, n.s.
	Residual	132	36.67		
Large fish	Ye	2	1.91	5.18	0.0068**
	Lo	1	0.74	4.03	0.0478*
	Si(Lo)	2	1.20	3.25	0.0402*
	Ye × Lo	2	1.99	5.40	0.0058**
	Ye × Si(Lo)	4	1.17	1.59	0.1810, n.s.
	Residual	132	24.36		

n.s., not significant.

Permutational univariate ANOVAs were based on Euclidean distances. P-values were obtained using 4999 permutations of residuals under a reduced model.

* $P < 0.05$.** $P < 0.01$.*** $P < 0.001$.

4. Discussion

4.1. Assessment of MPA impact

In general, results showed significant inside/outside differences in the multivariate abundance structure of fish assemblages, across years, for all groups of fish or species considered, except for unfished species. Multivariate interactions were also significant when environmental covariables were accounted for in the models. These results were confirmed and exemplified by significant univariate differences

between locations across years for total abundance, species richness and diversity for the abundances of the three species analysed.

Before MPA establishment, only the groups of small species, species of low fishing value and sedentary species already displayed significant inside/outside differences in abundance. Only species with low fishing value had on average higher abundances outside the MPA. The majority of sedentary species were small species (41%), whereas mobile species were mostly large species (39%). Thus, initial inside/outside differences could be explained by small species,

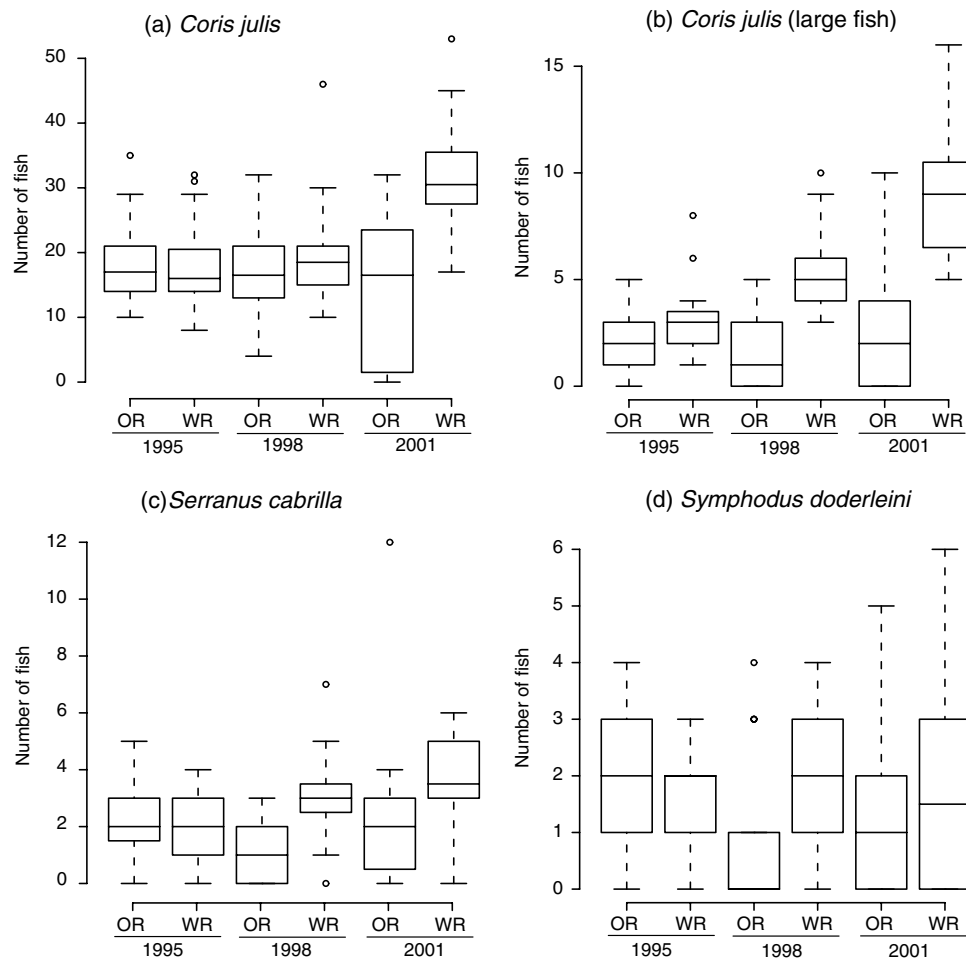


Fig. 4 – Boxplots of the abundances of (a) *Coris julis*, (b) large individuals of *Coris julis*, (c) *Serranus cabrilla*, and (d) *Symphodus doderleini* for each combination of the levels of factors Year and Location. There were $n = 24$ transects per combination.

with higher abundances inside the reserve. These species are usually sedentary with limited movements. Habitat preferences and/or natural variability could explain these spatial differences, even if habitat variables considered in the study did not allow to test this hypothesis. In 1995, indicator species could be identified only outside the MPA. The analysis of the fish assemblage status before MPA establishment is particularly important as it provides the baseline information for future monitoring and assessment (Edgar et al., 2004b).

After MPA establishment, changes in spatial patterns could be clearly evidenced (Fig. 2). In 1998, inside/outside differences in abundance were significant for all groups considered, except for small fish and surprisingly for large species. These corresponded to increased abundances within the MPA. The magnitude of the response to MPA establishment was not clearly related with fishing value at this early stage of restoration (i.e., three years after MPA establishment). At the fish assemblage level, inside/outside differences were more marked for metrics (total abundance, species richness and diversity) calculated from large fish only. At the species level, all metrics responded to MPA establishment, except for total abundance of *C. julis*, through increasing abundances within the MPA. From 1998 onwards, many species belonging to almost all the fami-

lies encountered in the study were significant indicator species within the MPA, but no indicator species could be identified for a particular year (1998 or 2001); which in fact would not be desirable for an indicator of protection.

It is interesting to note that, six years after MPA establishment (in 2001), inside/outside differences were even more significant than in 1998, except for metrics computed from small fish only. The contrast between increased abundances of large and medium fish, and stable abundances of small fish shows that six years after MPA establishment, positive effects mostly pertain to larger sizes and larger abundances within the MPA. Effects linked to reproduction are thus not yet evidenced, at least not from this kind of data. Furthermore, there is still no clear link between fishing value of species and response to MPA establishment. Although the differences in abundance of species with medium to high fishing value were more significant in 2001 than in 1998, this may rather be explained by demographic characteristics of species or changes in fishing patterns outside the MPA. This also explains why inside/outside differences became significant for *C. julis*, an important target species for fisheries. At the fish assemblage level, all metrics (total abundance, species richness and diversity) displayed significant inside/outside differences six years after MPA establishment. At this scale, metrics based on large

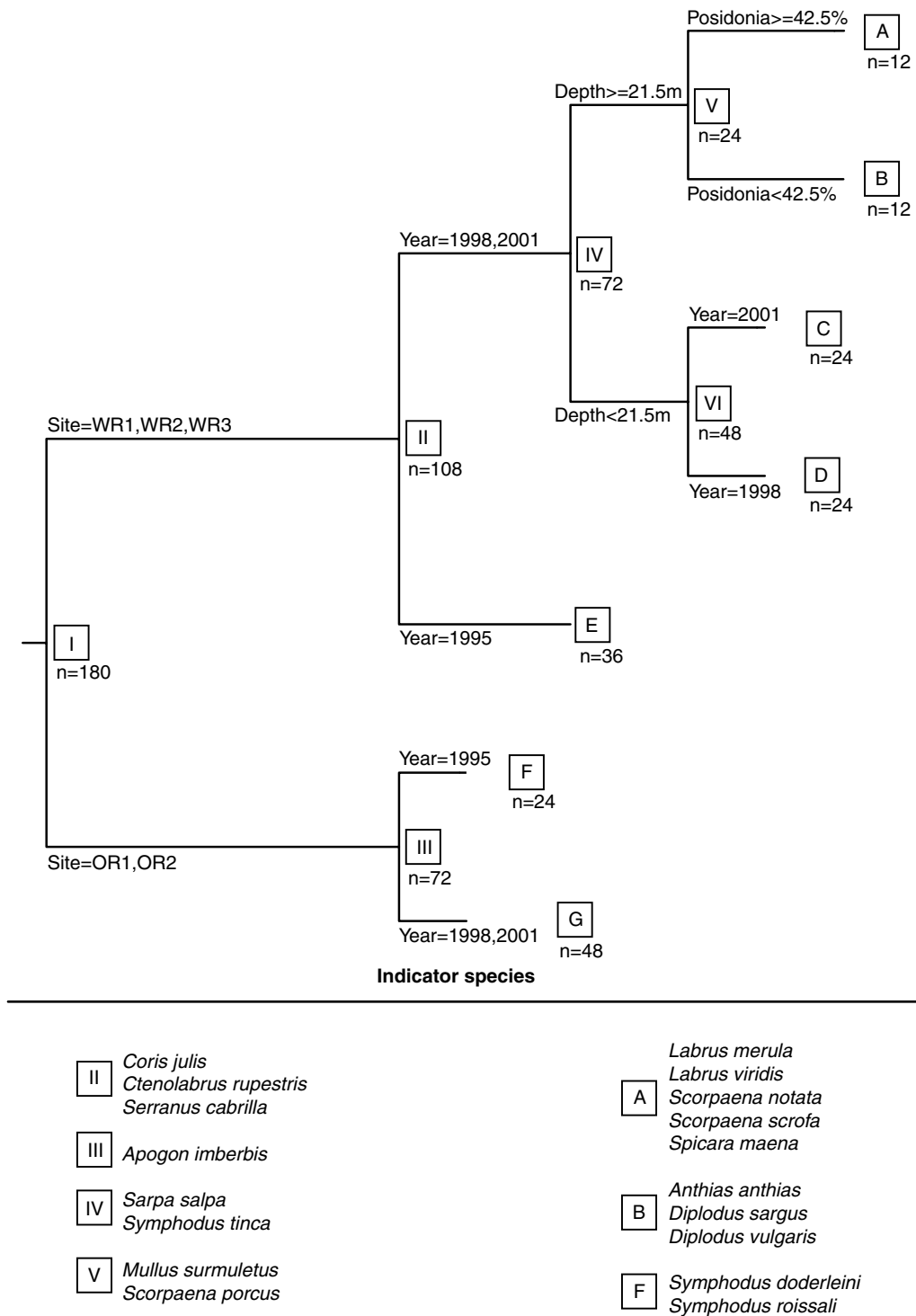


Fig. 5 – Multivariate regression tree constructed from the species abundance of large fish. For each split and each leaf, the number of transects was reported. Below the tree, indicator species for splits (in roman numbers) and leaves (in capitals) were reported.

fish thus appeared as relevant indicators of MPA effect at an early stage of restoration, while metrics calculated from all fish may be appropriate for restoration effects in the medium term.

Besides these effects of the Couronne MPA, mean depth and substrate complexity had an influence on the distribution of the whole fish assemblage and on large fish. Percentage lin-

ear cover of *P. oceanica* only had a structuring influence at depths deeper than 21.5 m and after 1995.

4.2. Methodological aspects

Studying MPA effectiveness by species or by taxa, as has been done in many articles, is important but not sufficient

in an ecosystem-based perspective. Changes in the composition of the whole fish assemblage have to be assessed across MPAs boundaries. The permutational multivariate analysis of variance allowed the production of a diagnostic on the evolution of the entire fish assemblage with respect to MPA establishment. Unlike MANOVA, the method does not require assumptions about distributions. The variance can be partitioned across the relevant factors of interest and any model can be tested. This analysis is rarely used to study MPA efficiency (Willis and Anderson, 2003; Fraschetti et al., 2005) (see comments on these studies below).

PERMANOVA informs about the effects of the MPA on the fish assemblage but it cannot be used directly to monitor the magnitude and direction of effects per metric. Ordination techniques, such as discriminant analysis, can be used to test and visualise these effects. MRT cannot be used for hypothesis testing but are helpful to build a hierarchical structure of the environmental variables structuring fish assemblages. Besides, it accepts quantitative variables and qualitative factors in the same analysis. MRT, together with IndVal, may provide indicators of MPA status. Note that the results obtained with the IndVal index were consistent and complementary with those obtained from the discriminant analysis.

In the case of persistent effects (“press impact”), as the effects caused by the establishment of an MPA on the fish assemblages are expected to be, the power of the statistical tests is not increased by an increase in the number of replicates, days or periods of sampling; but only by an increase in the number of locations of control and incidentally of impacted localities, if possible (Underwood and Chapman, 2003). Consequently, the experimental design used here to assess the effect of the MPA should be improved in the further monitoring programs by raising the number of locations, even if still taking habitat variables as covariables in the analyses.

4.3. Conservation aspects and indicators for management

The positive effects seen in this study may not be generalised. For example, in the Mimiwhangata Marine Park (New Zealand), the snapper (*Pagrus auratus*), the most heavily targeted fish species in the region, showed no difference in abundance or size between the Marine Park and adjacent control areas (Denny and Babcock, 2004). Fraschetti et al. (2005), studying benthic assemblages, showed that most of the variables considered (i.e., substrate cover, number of taxa, and average abundance of the most common taxa) were not significantly different between the protected and unprotected areas. Protecting species requires prioritisation. Indeed protection can improve abundances or sizes, but target species are very often predator species and thus there will be higher predation pressure on the preys inside the MPA, leading to changes in the fish assemblages (Francour, 1994; Pinnegar et al., 2000; Ashworth and Ormond, 2005). In the present study, notoriously cryptic fish species have been removed from the analyses, whereas MPAs could have potential negative impacts on them. Willis and Anderson (2003) showed that the sites inside the marine reserve contained, on average, lower densities of cryptic fishes than sites out-

side the reserve, which might be explained by effects of predators.

Defining groups of fish based on ecological or management criterion (e.g., commercial vs. non-commercial species) can provide different but complementary information about the status of the fish assemblage. The availability of data by size group is also helpful to assess the effects of protection, MPAs being effective at various temporal scales across fish sizes. Increase in fish size could be a direct effect of the protection and this increase could have indirect effects on inter-population differences in the reproductive output among organisms for which fecundity is dependent on body size (Roberts and Polunin, 1991). Actually, in a stochastic simulation model, including a subpopulation of larger mean, asymptotic body size resulted in less time spent at very small population sizes, which could reduce extinction risks (Kritzer and Davies, 2005).

An indicator is a metric that should be sensitive to the effect studied. Significant effects and their increased significance over time showed the sensitivity of the corresponding metrics to MPA establishment. From our results, relevant indicators could be at the species level, *S. cabrilla*, and to a lesser extent, *C. julis*, since they display marked differences only three years after MPA establishment. However, it is also interesting to monitor species that may display less immediate effects such as *S. doderleini*. At the fish assemblage level, metrics computed on large fish are obviously good indicators of restoration in early stages, whereas metrics computed on all fish give a more holistic appraisal at later stages.

Indicator species that are characteristic of a combination of factors and environmental variables such as depth may be a valuable tool for managers (Pullin et al., 2004). The more the habitat characteristics will be recorded precisely, the more accurate the index value of indicator species will be. The purpose here is not to avoid regular surveys identifying all the species encountered, but to be able to record, between such comprehensive surveys, the abundances of indicator species of particular interest for MPA monitoring. This monitoring of indicator species will inform about MPA efficiency at a reasonable cost, and it does not require costly training of observers. Such simplified monitoring protocols have already been investigated in the CBMP. They consisted of randomly spaced catching operations of large individuals of three Serranid species considered as indicators of a reserve effect (Harmelin et al., 1995).

Here, we assessed the effects of the MPA in relation to fisheries goals (i.e., effects on fish abundances and sizes) and conservation goals (i.e., effects on species richness and diversity). But many other aspects could have been investigated such as socio-economic impacts of the reserve (Badalamenti et al., 2000; Carter, 2003; Rudd et al., 2003; Chee, 2004; Pelletier et al., 2005). Linkages between ecological and economic systems often give rise to direct and immediate feedbacks (Brown et al., 2001). Clearly, studies on MPAs have to be more and more multidisciplinary, and this cannot be done without a clear planning, monitoring and evaluation (Jameson et al., 2002; Hilborn et al., 2004), and more linked with policy and management (Alder et al., 2002; Fazey et al., 2005).

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Appendix A

List of the 40 fish species recorded in the surveys with corresponding frequencies (in %) across the 3 years, for each site ($n = 36$ transects) and for each location ($n = 72$ for outside reserve (OR) and $n = 108$ for within reserve (WR)), and their class for each group analysed

Family	Species	Site					Location		Size	Fishing value	Mobility
		OR1	OR2	WR1	WR2	WR3	OR	WR			
Apogonidae	<i>Apogon imberbis</i>	2.8	16.7	2.8	2.8	0	9.7	1.9	1	U	M
Centracanthidae	<i>Spicara maena</i>	2.6	19.4	2.8	16.7	16.7	25	12	2	U	M
Centracanthidae	<i>Spicara smar</i>	5.6	2.8	2.8	0	11.1	4.2	4.6	1	U	M
Congridae	<i>Conger conger</i>	0	5.6	0	0	0	2.8	0	3	L	S
Labridae	<i>Coris julis</i>	100	83.3	100	100	91.7	91.7	97.2	2	MH	S
Labridae	<i>Ctenolabrus rupestris</i>	47.2	38.9	55.6	75	83.3	43.1	71.3	1	L	S
Labridae	<i>Labrus bimaculatus</i>	5.6	0	2.8	19.4	22.2	2.8	14.8	3	MH	S
Labridae	<i>Labrus merula</i>	16.7	22.2	11.1	33.3	19.4	19.4	21.3	3	MH	S
Labridae	<i>Labrus viridis</i>	11.1	11.1	5.6	8.3	8.3	11.1	7.4	3	MH	S
Labridae	<i>Symphodus cinereus</i>	0	5.6	0	0	0	2.8	0	1	MH	S
Labridae	<i>Symphodus doderleini</i>	77.8	52.8	80.6	72.2	47.2	65.3	66.7	1	L	S
Labridae	<i>Symphodus mediterraneus</i>	61.1	41.7	63.9	77.8	72.2	51.4	71.3	1	L	S
Labridae	<i>Symphodus melanocercus</i>	52.8	50	77.8	72.2	69.4	51.4	73.1	1	L	S
Labridae	<i>Symphodus ocellatus</i>	2.6	22.2	33.3	19.4	2.8	26.4	18.5	1	L	S
Labridae	<i>Symphodus roissali</i>	25	33.3	27.8	16.7	0	29.2	14.8	1	L	S
Labridae	<i>Symphodus rostratus</i>	2.6	36.1	36.1	2.6	13.9	33.3	26.9	1	L	S
Labridae	<i>Symphodus tinca</i>	38.9	2.6	47.2	41.7	19.4	34.7	36.1	2	MH	S
Moronidae	<i>Dicentrarchus labrax</i>	0	0	5.6	0	0	0	1.9	3	MH	M
Mugilidae	<i>Chelon labrosus</i>	0	0	2.8	0	0	0	0.9	3	L	M
Mullidae	<i>Mullus surmuletus</i>	27.8	19.4	44.4	27.8	2.6	23.6	34.3	2	MH	M
Muraenidae	<i>Muraena helena</i>	0	5.6	0	0	0	2.8	0	3	L	S
Pomacentridae	<i>Chromis chromis</i>	86.1	77.8	86.1	83.3	94.4	81.9	88	1	U	M
Scorpaenidae	<i>Scorpaena notata</i>	0	0	2.8	2.8	8.3	0	4.6	1	MH	S
Scorpaenidae	<i>Scorpaena porcus</i>	0	5.6	0	8.3	27.8	2.8	12	2	MH	S
Scorpaenidae	<i>Scorpaena scrofa</i>	2.8	0	0	2.8	11.1	1.4	4.6	3	MH	S
Serranidae	<i>Anthias anthias</i>	0	0	0	0	5.6	0	1.9	1	U	M
Serranidae	<i>Epinephelus caninus</i>	0	0	0	0	2.8	0	0.9	3	MH	S
Serranidae	<i>Epinephelus marginatus</i>	0	0	0	0	5.6	0	1.9	3	MH	S
Serranidae	<i>Serranus cabrilla</i>	97.2	61.1	94.4	97.2	75	79.2	88.9	2	MH	S
Serranidae	<i>Serranus scriba</i>	13.9	11.1	2.8	2.8	0	12.5	1.9	2	MH	S
Sparidae	<i>Boops boops</i>	47.2	11.1	11.1	16.7	33.3	29.2	1.4	2	L	M
Sparidae	<i>Dentex dentex</i>	0	0	2.8	0	2.8	0	1.9	3	MH	M
Sparidae	<i>Diplodus annularis</i>	2.8	8.3	2.8	5.6	0	5.6	2.8	1	MH	M
Sparidae	<i>Diplodus sargus</i>	22.2	16.7	36.1	58.3	44.4	19.4	46.3	2	MH	M
Sparidae	<i>Diplodus vulgaris</i>	16.7	38.9	33.3	41.7	52.8	27.8	42.6	2	MH	M
Sparidae	<i>Oblada melanura</i>	0	0	2.8	0	0	0	0.9	1	L	M
Sparidae	<i>Pagellus erythrinus</i>	0	0	0	0	2.8	0	0.9	3	MH	M
Sparidae	<i>Sarpa salpa</i>	5.6	13.9	19.4	5.6	8.3	9.7	11.1	3	L	M
Sparidae	<i>Sparus pagurus</i>	2.8	0	2.8	2.8	2.8	1.4	2.8	3	MH	M
Sparidae	<i>Spondyliosoma cantharus</i>	0	2.8	2.8	0	2.8	1.4	1.9	3	MH	M

Size groups were defined by species of minimum and maximum length within the range 8–20 cm (1), 20–30 cm (2), and 30–200 cm (3). Fishing value groups corresponded to unfished species (U) and species with low (L) or medium to high (MH) fishing value. Mobility groups were defined by sedentary (S) or mobile (M) species.

Appendix B

Mean species abundances per site and per year (\pm SE)

Group/species considered	1995		1998		2001	
	WR	OR	WR	OR	WR	OR
<i>Fish sizes</i>						
All	3.24 \pm 0.73	2.58 \pm 0.49	2.88 \pm 0.61	2.46 \pm 0.62	6.3 \pm 3.1	2.32 \pm 0.58
Large	1.22 \pm 0.58	0.65 \pm 0.24	0.74 \pm 0.17	0.45 \pm 0.23	0.56 \pm 0.08	0.56 \pm 0.32
Medium	1.8 \pm 0.43	1.25 \pm 0.32	1.8 \pm 0.55	1.89 \pm 0.56	5.51 \pm 3.1	1.68 \pm 0.44
Small	0.21 \pm 0.06	0.68 \pm 0.25	0.34 \pm 0.1	0.11 \pm 0.03	0.22 \pm 0.05	0.08 \pm 0.02
<i>Species sizes</i>						
Large	0.01 \pm 0.01	0.02 \pm 0.01	0.15 \pm 0.08	0.08 \pm 0.02	0.27 \pm 0.13	0.02 \pm 0.01
Medium	2.66 \pm 0.39	2.81 \pm 0.55	2.64 \pm 0.41	3.02 \pm 0.44	4.71 \pm 0.67	3.68 \pm 0.97
Small	7.28 \pm 2.01	5.31 \pm 1.3	6.15 \pm 1.68	4.77 \pm 1.71	14.21 \pm 8.67	4.03 \pm 1.48
<i>Fishing value</i>						
Unfished	20.23 \pm 5.86	14.35 \pm 14.35	17.16 \pm 4.91	14.77 \pm 5.06	41.35 \pm 25.9	12.49 \pm 4.38
Low	0.78 \pm 0.1	0.98 \pm 0.33	0.63 \pm 0.11	0.65 \pm 0.11	1.07 \pm 0.25	1.36 \pm 0.68
Medium to high	1.03 \pm 0.16	0.99 \pm 0.16	1.13 \pm 0.18	0.87 \pm 0.15	1.84 \pm 0.28	0.73 \pm 0.73
<i>Species mobility</i>						
Mobile species	5.58 \pm 1.59	4.13 \pm 1.05	4.78 \pm 1.34	4.17 \pm 1.36	11.58 \pm 6.85	4.09 \pm 1.26
Sedentary species	1.3 \pm 0.17	1.29 \pm 0.17	1.31 \pm 0.18	1.04 \pm 0.15	1.94 \pm 0.28	0.86 \pm 0.16
<i>Community metrics</i>						
Total fish abundance	135.92 \pm 23.40	108.21 \pm 14.12	121.10 \pm 18.89	103.12 \pm 22.54	264.71 \pm 127.56	97.58 \pm 23.97
Total fish abundance (L)	51.33 \pm 23.41	27.17 \pm 9.73	31.12 \pm 6.77	18.80 \pm 9.94	23.71 \pm 3.27	23.71 \pm 13.48
Species richness	8.71 \pm 0.57	9.92 \pm 0.51	9.10 \pm 0.43	9.04 \pm 0.53	9.87 \pm 0.60	5.12 \pm 0.78
Species richness (L)	5.21 \pm 0.44	5.83 \pm 0.47	6.87 \pm 0.47	4.04 \pm 0.55	6.5 \pm 0.61	2.46 \pm 0.37
Diversity	1.53 \pm 0.13	1.70 \pm 0.11	1.63 \pm 0.12	1.78 \pm 0.12	1.82 \pm 0.14	1.05 \pm 0.16
Diversity (L)	1.65 \pm 0.17	1.86 \pm 0.18	2.12 \pm 0.13	1.40 \pm 0.17	1.91 \pm 0.13	0.83 \pm 0.14
<i>Fish species</i>						
<i>Coris julis</i>	18.00 \pm 1.35	18.00 \pm 1.25	19.6 \pm 1.52	16.70 \pm 1.21	31.67 \pm 1.65	14.33 \pm 2.25
<i>Coris julis</i> (L)	2.92 \pm 0.33	2.46 \pm 0.29	5.37 \pm 0.41	1.62 \pm 0.33	9.10 \pm 0.61	2.54 \pm 0.54
<i>Serranus cabrilla</i>	2.17 \pm 0.21	2.29 \pm 0.28	3.12 \pm 0.30	1.08 \pm 0.18	3.71 \pm 0.34	2.21 \pm 0.50
<i>Serranus cabrilla</i> (L)	0.33 \pm 0.11	0.62 \pm 0.18	2.46 \pm 0.24	0.29 \pm 0.13	1.54 \pm 0.24	0.17 \pm 0.10
<i>Symphodus doderleini</i>	1.58 \pm 0.21	1.92 \pm 0.20	1.75 \pm 0.23	0.75 \pm 0.23	1.83 \pm 0.37	1.04 \pm 0.26
<i>Symphodus doderleini</i> (L)	0.96 \pm 0.16	1.17 \pm 0.18	1.37 \pm 0.21	0.75 \pm 0.16	0.67 \pm 0.18	0.50 \pm 0.13
Site WR3 removed from WR location. Species with null abundance in a group were kept in the mean and standard error computations. (L) means large fish.						

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